Molecular Electrocatalysis for Oxygen Reduction by Cobalt Porphyrins Adsorbed at Liquid/Liquid Interfaces

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Abstract: Molecular electrocatalysis for oxygen reduction at a polarized water/1,2-dichloroethane (DCE) interface was studied, involving aqueous protons, ferrocene (Fc) in DCE and amphiphilic cobalt porphyrin catalysts adsorbed at the interface. The catalyst, (2,8,13,17-tetraethyl-3,7,12,18-tetramethyl-5-aminophenylporphyrin) cobalt(II) (CoAP), functions like conventional cobalt porphyrins, activating O2 via coordination by the formation of a superoxide structure. Furthermore, due to the hydrophilic nature of the aminophenyl group, CoAP has a strong affinity for the water/DCE interface as evidenced by lipophilicity mapping calculations and surface tension measurements, facilitating the protonation of the CoAP–O2 complex and its reduction by ferrocene. The reaction is electrocatalytic as its rate depends on the applied Galvani potential difference between the two phases.

Introduction

Molecular oxygen (O2) reduction reaction (ORR) is an important topic in biological and energy-related chemistry. However, it is a spin-forbidden process, which is kinetically slow at ambient temperature unless a catalyst is present. Nature has made the choice in aerobic organisms by evolving membrane-bound enzymes that contain a porphyrin substructure as catalysts. Inspired by their role in nature, a number of metalloporphyrins have been chemically synthesized. Their catalytic activity has been investigated either electrochemically using the modified electrode methodology or chemically using molecular electron donors, such as ferrocene (Fc) and its derivatives. The Porphyrin Handbook, Kadish, K. M., Smith, K. M., Guilard, R., Eds.; Academic Press: San Diego 2003; Vol. 11, pp 1–49.

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derivatives, in homogeneous solutions.\textsuperscript{26–28} In the latter case, reduction of O\textsubscript{2} by Fe in acidic solutions\textsuperscript{29–32} proceeds rather slowly, and the presence of a catalytic amount of metalloporphyrins can significantly accelerate the reaction rate.\textsuperscript{26–28}

On the other hand, ORR is a proton-coupled electron transfer (PCET) reaction. When studying ORR on a solid electrode by amperometry, one measures the electron transfer rate, but it is not possible to control the proton transfer step by controlling the electrode potential. Electrochemistry at a liquid/liquid interface, also called interface between two immiscible electrolyte solutions (ITIES), has manifested itself recently as a unique approach to study PCET reaction with the possibility of locating protons in the aqueous phase and electron donors in the organic phase.\textsuperscript{33} In this work, proton-coupled oxygen reduction by Fe involving an amphiphilic cobalt porphyrin catalyst, (2,8,13,17-tetraethyl-3,7,12,18-tetramethyl-5-p-aminophenylporphyrin) cobalt(II) (CoAP), at a polarized water/1,2-dichloroethane (DCE) interface is reported. CoAP serves as a redox catalyst like conventional cobalt porphyrins, activating O\textsubscript{2} reduction via coordination with the cobalt(II) (Co\textsuperscript{III}) center by the formation of a superoxide structure. Moreover, since CoAP is an amphiphilic molecule with a strong interfacial affinity, the proton-coupled oxygen reduction is an interfacial process. These two factors play together to account for the excellent electrocatalytic activity of CoAP, which is better than those previously reported for cobalt porphine (CoP),\textsuperscript{33} cobalt tetraphenylporphyrin (CoTPP),\textsuperscript{34} and cobalt octaethylporphyrin Co(OEP)\textsuperscript{35} that are not amphiphilic. The present system provides an example of interfacial molecular electrocatalysis for oxygen reduction combining the advantages of molecular catalyst and electrocatalysis because the ORR depends on the applied potential.

Experimental Section

Chemicals. All chemicals were used as received without further purification. The aqueous solutions were prepared with ultrapure water (18.2 M\textsubscript{2} cm\textsuperscript{-1}). Ferrocene (Fe, 98%), 1,1′-dimethylferrocene (DFc, 97%), ferrocenecarboxaldehyde (FcCA, 98%), trifluoroacetic acid (TFA, 98%), and lithium tetrakis(pentafluorophenyl)borate (LiTB, diethyl etherate were purchased from Aldrich. Lithium acid (TFA, 98%), and lithium tetrakis(pentafluorophenyl)borate (DFc, 97%), ferrocenecarboxaldehyde (FcCA, 98%), trifluoroacetic acid (HCl, 32%) was bought from Merck. Bis(triphenylphosphine) octyldiene)ammonium tetrakis(pentafluorophenyl)borate (BATB) was prepared as previously reported.\textsuperscript{36}

Surface Tension Measurements. Surface tension measurements were performed using the video-image pendant-drop method in a four-electrode all-glass cell, which was adapted from a spectrosocpic cuvette (10 mm × 10 mm). The cell was connected to a four-electrode potentiostat (PGSTAT 30, Eco-Chemie, Netherlands). The cell potential was changed stepwise (25 mV per step, 25 s step time), and on each step the drop image was captured (24 s after the potential step change), and analyzed with the help of CAM101 System (KSV Instruments Ltd., Finland). Potential scale conversion was based on the reversible half-wave potential of the TMA\textsuperscript{+} ion transfer (0.160 V).\textsuperscript{37} Measurements were carried out under aerobic conditions unless specified otherwise.

Two-Phase Reactions Controlled by the Distribution of a Common Ion. Two-phase reactions were performed in small glass flasks with a volume of 10 mL. A flask was filled first with 2 mL of DCE solution containing reactants (Fe, CoAP), followed by the addition of 2 mL of aqueous solution containing 10 mM HCl. The salts of the common ion (TB\textsuperscript{+}), LiTB and BATB, were added in the same concentration of 5 mM to the aqueous and DCE phases, respectively. After stirring and further waiting for the clear separation of two phases, the aqueous and organic solutions were isolated from each other. The organic phase was directly subjected to the UV-visible spectrophotometric measurement (Ocean Optical CHEM2000 spectrophotometer, quartz cuvette with a path length 10 mm), while the aqueous phase was first treated by excess NaI (equivalent to 0.1 M) prior to the UV-visible spectrophotometric measurement.

Lipophilicity Mapping. The lipophilicity distribution around molecules was characterized by the molecular lipophiliity potential

\textsuperscript{(38) Girault, H. H.; Schiffrin, D. J.; Smith, B. D. V. J. Colloid Interface Sci. 1984, 101, 257–266.}
(MLP) calculated at any given point in space around a molecule using two components for the calculation, namely a fragmental system of lipophilicity and a distance function as expressed by eq 1:

$$MLP_k = \sum_{i=1}^{N} f_i \cdot fct(d_{ik})$$

where $k$ indicates a given point in space, $i$ a given molecular fragment, $N$ the total number of fragments in the molecule, $f_i$ the lipophilic increment of fragment $i$, $fct$ a distance function, and $d_{ik}$ the distance between fragment $i$ and point $k$.

Two different approaches were used to calculate the MLP, namely the CLIP software\(^{40,41}\) and MolCAD software\(^{42}\) as implemented in SYBYL 8.0.\(^{43}\) CLIP calculates the MLP on the water accessible surface area of a compound and displays the result as colored dots, whereas the MOLCAD calculates the MLP on its molecular surface and displays the result as a colored surface. The two approaches used different atomic fragmental systems, namely the Broto and Moreau system\(^ {44}\) for CLIP and the Ghose and Crippen system\(^ {45,46}\) for MolCAD but a similar distance function reported in eq 2:

$$MLP_k = \sum_{i=1}^{N} f_i \cdot \left(1 + \frac{e^{-a'b}}{1 + e^{-c'(d_{ik}-b)}}\right)$$

where $a$ and $b$ define respectively the decay and the distance cutoff for the sigmoidal variation of the MLP with respect to the distance ($d_{ik}$) between the dot $k$ and the fragment $i$. A fragmental value has to be assigned to the cobalt atom not parametrized in both fragmental atomic systems. The qualitative distribution of lipophilicity on molecular surfaces was not largely affected by large variations of the fragmental values for the cobalt atom defined as a polar monocentric atom. Thus, a fragmental value calibrated using atomic charges was retained for the cobalt atom.

In CLIP representations, the color coding follows a scale starting from the most polar to the most hydrophobic regions, namely red, orange, yellow, white, green, green-blue, blue. In MolCAD representations, the color coding follows a scale starting from the most polar to the most hydrophobic regions, namely blue, green-blue, green, brown, dark brown.

### Results and Discussion

**Synthesis of CoAP.** The synthetic pathway is given in Scheme 2. First, 2,8,13,17-tetraethyl-3,7,12,18-tetramethyl-5-$p$-aminophenylporphyrin (H$_2$AP) was synthesized by a method different from that described previously by Senge et al.\(^ {47}\) Indeed, in this previous report, the authors were reacting H$_2$OEP (OEP: 2,3,7,8,12,13,17,18-octapyrlylopyrphenylporphyrin) with an excess of the organolithium reagent of $p$-bromoaniline at low temperature. In the present case, the starting material was not the corresponding free base but the $1,19$-dideoxy-3,8,12,17-tetraethyl-2,7,13,18-tetramethylbiladien-$a,c$-dibromide already described by our group and readily available in large amounts.\(^ {48}\) Reaction of $p$-nitrobenzaldehyde with the $a,c$-biladien gave the corresponding nitroporphyrin which was subsequently converted to H$_2$AP by reduction with tin(II) dichloride. CoAP was obtained by reacting H$_2$AP with cobalt acetate tetrahydrate in a refluxing CHCl$_3$/MeOH mixture. All compounds were fully characterized by HRMS, $^{1}$H NMR, and UV–visible spectroscopy. Detailed synthetic procedures and spectroscopic data are given in the Supporting Information.

**Interfacial Adsorption and Protonation of CoAP.** CVs in the absence and presence of 50 μM CoAP in DCE are shown in Figure 1a. The current observed at the negative and positive ends of the potential window is due to the transfer of Cl$^-$ and H$^+$ from water to DCE, respectively. Compared to the background current, the presence of CoAP resulted in a large capacitive current indicating the presence of an adsorbed layer. To verify this point, surface tension measurements were

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performed using a video-image pendant-drop method, and the obtained electrocapillary curves are displayed in Figure 1b. The parabolic shape of the electrocapillary curve in the absence of CoAP is characteristic of two back-to-back Gouy–Chapman diffuse layers without any specific adsorption of electrolyte ions. The addition of CoAP in DCE results in a significant decrease in the surface tension primarily at potentials negative to the electrocapillary maximum (potential of zero charge, \( \text{pzc} \)), indicating the adsorption of CoAP species. Toward positive potentials, the surface tension starts to increase and finally overlapped with the surface tension of the background electrolytes. It has to be mentioned that the measurements started from negative potentials, but that surface tension data obtained in the reverse direction (blue) is the same.

The adsorption of CoAP stems from its amphiphility as illustrated in Figure 2, with the amino group as the hydrophilic moiety and the metalated ring as the lipophilic part. The interfacial activity of CoAP is also related to its acid–base properties and to the concentration of protons on the aqueous side of the interface, which increases at potentials positive to the \( \text{pzc} \). Thus, the significant decrease of the surface tension at very negative potentials can be attributed to the adsorption of neutral CoAP and oxygenated CoAP, \((\text{Co}–\text{O}_2)\text{AP}\), at the interface. As shown below, CoAP and \((\text{Co}–\text{O}_2)\text{AP}\) coexist in the stock sample both in solid and in solution. When scanning the potential more positive than the \( \text{pzc} \), the proton concentration at the interface is increased and the adsorbed \((\text{Co}–\text{O}_2)\text{AP}\) can be protonated to form \([\text{(Co}–\text{O}_2\text{H})\text{AP}]^+\) by protonation of the bound oxygen. The protonation of the amino group usually takes place at much more positive potentials or at much lower pH values. The adsorbed \([\text{(Co}–\text{O}_2\text{H})\text{AP}]^+\) will desorb to the organic phase at a positive potential, leading to a decrease of interfacial coverage of \([\text{(Co}–\text{O}_2\text{H})\text{AP}]^+\) and an increase of the surface tension. The surface tension data thereby point to the formation of \([\text{(Co}–\text{O}_2\text{H})\text{AP}]^+\) that undergoes an electrochemical ion transfer reaction from the adsorbed state:

\[
\begin{align*}
\text{(Co}–\text{O}_2)\text{AP}^{\text{(ads)}} + \text{H}^+_\text{(w)} & \rightarrow [\text{(Co}–\text{O}_2\text{H})\text{AP}]^{\text{(ads)}}_+ \quad (3) \\
[\text{(Co}–\text{O}_2\text{H})\text{AP}]^{\text{(ads)}}_+ & \rightarrow [\text{(Co}–\text{O}_2\text{H})\text{AP}]^{\text{(DCE)}}_+ \quad (4)
\end{align*}
\]

Here, the species in water, in DCE and adsorbed at the interface are denoted as (w), (DCE), and (ads), respectively. It is well-known that cobalt porphyrins are able to coordinate O\(_2\), thereby forming a superoxide-like structure, which can be protonated to produce hydroperoxy radicals.

Figure 3a shows that the surface tension decreases considerably with increasing the concentration of CoAP in DCE. This dependence can be used to evaluate the surface excess concentration of CoAP and to determine the adsorption isotherm on the interface.


Activation of ORR by CoAP at a Water/DCE Interface

Figure 3. (a) Electrocapillary curves at various concentrations of CoAP using Cell 2: z = 10 μM (black), 25 μM (red), 50 μM (blue), 100 μM (green), and 200 μM (brown). The measurements started from the negative potential. (b) CoAP adsorption isotherm constructed with the surface tension data at −0.10 V. The red line corresponds to a fit to the Langmuir isotherm.

The basis of the Gibbs adsorption equation (see the derivation in the Supporting Information):53

\[
\Gamma_{\text{CoAP}} + \Gamma_{(\text{Co}-\text{O}_2)\text{AP}} + \Gamma_{[(\text{Co}-\text{O}_2\text{H})\text{AP}]^+} = \frac{-1}{RT} \left( \frac{\partial \gamma}{\partial \ln c_{\text{CoAP}}} \right)_{T,P,\Delta\Phi,\mu_{\text{CoAP}}} \tag{5}
\]

where \(c_{\text{CoAP}}\) is the bulk concentration of CoAP in DCE and \(\Gamma_i\) (\(i = \text{CoAP}, (\text{Co}-\text{O}_2)\text{AP}\) and \([(\text{Co}-\text{O}_2\text{H})\text{AP}]^+)\) represent the respective surface excess concentration. In terms of eq 5, the surface coverage of CoAP species has three contributions. At positive potentials \([(\text{Co}-\text{O}_2\text{H})\text{AP}]^+\) is formed, the deconvolution of this contribution from two others is difficult because both the adsorption and transfer processes are involved. However, at potentials more negative than −0.1 V, the adsorption of neutral CoAP and (Co−O2)AP is dominant. Assuming, for simplicity, that CoAP and (Co−O2)AP have the same molecular structure and interfacial affinity, the concentration dependence of the surface excess concentration evaluated at −0.1 V (shown in Figure 3b), can be fitted to a Langmuir isotherm:

\[
\Gamma_{\text{CoAP}} = \frac{\beta_{\text{CoAP}}}{1 + \beta_{\text{CoAP}} c_{\text{CoAP}}} \tag{6}
\]

where \(\Gamma_{\text{CoAP}}\) and \(\beta\) are the maximum surface excess concentration and Langmuir adsorption coefficient of CoAP, respectively.

The fitting yields \(\Gamma_{\text{CoAP}}^* = (1.03 \pm 0.04) \times 10^{-6} \text{ mol m}^{-2}\) and \(\beta = (3.29 \pm 0.04) \times 10^4 \text{ M}^{-1}\). From \(\Gamma_{\text{CoAP}}^*\), the area per molecule occupied amounts 161 Å2, which suggests that adsorbed CoAP species take an orientation normal to the interface with the hydrophilic amino group toward the aqueous phase.

Interfacial Proton-Coupled Oxygen Reduction Catalyzed by CoAP. Figure 4a compares the CVs obtained under different experimental conditions. First, in the presence of only Fc in DCE under aerobic conditions a small voltammetric wave with a half-wave potential at \(\Delta_{w}\Phi_{1/2} = 0 \text{ V}\) was observed, which corresponds to the transfer of ferrocenium (Fc\(^+\)) produced by a slow oxidation of Fc in air of the stock solution.33,34 Also shown above in Figure 1a is that in the presence of only CoAP in DCE under aerobic conditions only an overall capacitive current signal was observed. However, when both Fc and CoAP were dissolved in DCE, an irreversible positive current signal appeared in the positive potential range. A control experiment under anaerobic conditions showed that this current signal did not appear (blue line in Figure 4a). These facts suggest that CoAP, O2, and Fc must be present at the same time to observe this signal. Therefore, similar to that observed with cobalt porphine,33 this irreversible current signal corresponds to a proton-coupled oxygen reduction process catalyzed by CoAP. This process produces Fe\(^+\), thus accounting for the significant increment of the Fe\(^+\) transfer current wave located at 0 V as shown in Figure 4a. Indeed, the standard redox potential for Co\(^{3+}\text{AP}/\text{Co}^{2+}\text{AP}\) is equal to 0.69 V vs SHE (see Supporting Information, Figure S-1) more positive than that of Fe\(^+\) equal to 0.64 V vs SHE. In

fact, when cycling the potential to the positive values repeatedly, more Fc\(^+\) will be produced, leading to a continuous increase of the Fc\(^+\) transfer current scan after scan (black lines in Figure 4b). In contrast, under anaerobic conditions, the Fc\(^+\) transfer current wave did not increase with repeated potential scans (red lines in Figure 4b).

The reduction of O\(_2\) by Fc was also investigated by so-called shake-flask experiments performed as reported previously.\(^{33}\) The Galvani potential difference across the water/DCE interface was polarized at a very positive value of 0.54 V by two salts, LiTB and BATB, having a common anion of TB\(^-\). At such potential values, protons are transferred from water to DCE by the lipophilic aqueous TB\(^-\) ions acting as a proton pump dragging with them protons as they transfer to the organic phase. Results of the shake-flask experiments are shown in Figure 5. As shown in Figure 5a, 10 mM HCl was present in water in all three flasks, and the DCE phase contained only 1 mM Fc in flask 1, only 20 \(\mu\)M CoAP in flask 3, and both in flask 2. It was observed that the DCE phase in flask 2 changed its color from pink (inset in (a)) to dark green immediately after being put in contact with the water solution. In contrast, the DCE phase in the other two flasks remained the same. The two phases were then separated from each other for further spectroscopic and colorimetric tests.

First, UV–visible spectra of the separated DCE solutions were measured, as shown in Figure 6a. The formation of Fc\(^+\) in the DCE solution from flask 2 was revealed, and the absorption band with a maximum at 620 nm represents its signature (blue line). The color change is thus due to the oxidation of Fc to Fc\(^+\), and the color of DCE phase in the flask 2 reflects a mixed color of green Fc\(^+\) and pink CoAP. In contrast, in the presence of Fc only in DCE (flask 1) no Fc\(^+\) was detected, because no changes in color and in the absorption spectrum (red line compared to full black line) were observed. On the other hand, when only CoAP is present in DCE (flask 3), changes in the absorption spectrum (green line compared to dotted black line) due to the oxidation of CoIIAP to CoIIIAP\(^+\) and the protonation of the amino group (see Supporting Information, Figure S-2) were observed. As for the isolated aqueous solutions from three flasks, excess NaI equivalent to 0.1 M was added. It was observed that the color from flask 2 changed from colorless to yellow as illustrated in Figure 5b. As shown previously, this color change can be due to the presence of H\(_2\)O\(_2\) in the solution.\(^{54}\) H\(_2\)O\(_2\) is a strong oxidant that can oxidize I\(^-\) to I\(_3^--\), which appears yellow and displays two absorption bands (\(\lambda_{max} = 286\) and 352 nm) as shown by

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the blue line in Figure 6b. Additionally, I$_3^-$ changes its color to very brown when meeting with aqueous soluble starch (Figure 5c). In contrast, no H$_2$O$_2$ was detected at all in the aqueous solutions from flasks 1 and 3, as seen both from the colorimetric responses and the UV–visible spectra (green curve in Figure 6b).

The above experimental facts clearly demonstrated that H$_2$O$_2$ and Fc$^+$ were produced in water and DCE, respectively, only when both Fc and CoAP were present (flask 2). A control experiment under anaerobic conditions (Figure 5d) showing that the DCE solution containing Fc and CoAP did not change its color even after 3 h, further proving the reduction of O$_2$ to H$_2$O$_2$ by Fc catalyzed by CoAP. Similar to other conventional cobalt porphyrins, CoAP activates O$_2$ by the formation of a superoxide-like intermediate with O$_2$ coordinated to cobalt, which can be seen clearly in the UV–visible spectrum of CoAP. As shown in Figure 6a, a freshly prepared CoAP solution displays two Soret bands ($\lambda_{\text{max}}$ = 397 and 423 nm) and a broad convoluted Q-band ($\lambda_{\text{max}}$ = 552 nm). The second Soret band reflects that the solution contains (Co−O$_2$)AP in which the electron delocalization from Co(II) to O$_2$ occurs, leading to a bathochromic shift of both Soret and Q bands in the UV–visible spectrum.$^{51,52}$

**Mechanism and Thermodynamic Analysis.** An overall reaction of oxygen reduction with Fc catalyzed by CoAP is:

$$2\text{Fc}^{+\text{(DCE)}} + 2\text{H}^+_{(W)} + \text{O}_2^{\text{(DCE)}} \rightarrow 2\text{Fc}^{+\text{(DCE)}} + \text{H}_2\text{O}_2(\text{W})$$

(7)

A corresponding reaction scheme involving the adsorption of (Co−O$_2$)AP (dashed line), proton-coupled oxygen reduction (solid line), and regeneration of CoAP$^+$ by Fc (dotted line) is illustrated in Scheme 3. These steps can be expressed as follows:

$$\text{CoAP}^{\text{(DCE)}} + \text{O}_2^{\text{(DCE)}} \rightarrow (\text{Co−O}_2)\text{AP}^{\text{(DCE)}}$$

(8)

$$\text{(Co−O}_2)\text{AP}^{\text{(DCE)}} \rightarrow (\text{Co−O}_2)\text{AP}^{\text{(ads)}}$$

(9)

$$\text{(Co−O}_2)\text{AP}^{\text{(ads)}} + 2\text{H}^+_{(W)} + \text{Fc}^{\text{(DCE)}} \rightarrow \text{CoAP}^{\text{+\text{(DCE)}}} + \text{Fc}^{\text{+\text{(DCE)}}} + \text{H}_2\text{O}_2(\text{W})$$

(10)

$$\text{CoAP}^{\text{+\text{(DCE)}}} + \text{Fc}^{\text{(DCE)}} \rightarrow \text{CoAP}^{\text{(DCE)}} + \text{Fc}^{\text{+\text{(DCE)}}}$$

(11)

In terms of this scheme, the Nernst equation for the overall process is:

$$\Delta_{\text{w}}^\circ \phi_{\text{el}} = \Delta_{\text{w}}^\circ \phi_{\text{el}}^0 + \frac{RT \ln 10}{F} \text{pH} +$$

$$\frac{RT}{2F} \ln \left( \frac{a_{\text{FC}^+}}{a_{\text{FC}^0}} \cdot \frac{a_{\text{CoAP}^+}}{a_{\text{CoAP}^0}} \cdot \frac{p_{\text{O}_2}}{a_{\text{H}_2\text{O}_2}^w} \right)$$

(12)

where $a_{\alpha}^i (\alpha = o$ or $w$) represents the activity of species $i$, $p^0$ and $f_{o_2}$ are the standard pressure and the fugacity of oxygen. $\Delta_{\text{w}}^\circ \phi_{\text{el}}^0$ represents the standard Galvani potential difference given by:

$$\Delta_{\text{w}}^\circ \phi_{\text{el}}^0 = \frac{1}{2} \left[ \frac{E_{\text{CoAP}^{\text{+\text{(DCE)}}}/\text{CoAP}^{\text{(DCE)}}}}{\text{SHE}} + \frac{1}{2} \left( \frac{E_{\text{FC}^{\text{+\text{(DCE)}}}/\text{FC}^{\text{(DCE)}}}}{\text{SHE}} - \frac{E_{\text{O}_2^{\text{(DCE)}}}/\text{H}_2\text{O}_2}}{\text{SHE}} \right) \right]$$

(13)

where $E_{\text{CoAP}^{\text{+\text{(DCE)}}}/\text{CoAP}^{\text{(DCE)}}}$, $E_{\text{FC}^{\text{+\text{(DCE)}}}/\text{FC}^{\text{(DCE)}}}$, and $E_{\text{O}_2^{\text{(DCE)}}}/\text{H}_2\text{O}_2}$ denote the standard redox potentials of the CoAP$^+/CoAP$ and Fc$^+/Fc$ couples in DCE and the O$_2/H_2O_2$ couple in water, respectively, all with respect to the aqueous standard hydrogen electrode (SHE). Equation 12 predicts the pH dependence of the redox Galvani potential difference, which is corroborated by the shift of the current signal with pH by about 60 mV/pH as shown in Figure 7. Equations 12 and 13 also indicate a dependence on the standard redox potential of the organic electron donor. When Fc is replaced either by a stronger electron donor, DFc, or by a weaker one, FcCA, the onset potential of the irreversible current shifts to negative and positive potentials, respectively, as displayed in Figure 8a. A negative shift of 45 mV was experimentally observed for DFc with a logarithmic plot (Figure
8b), which is in agreement with the theoretical prediction of eqs 12 and 13.

Finally, it should be mentioned that the proton-coupled oxygen reduction catalyzed by CoAP is an interfacial process, as voltammetry only measures interfacial charge transfer reactions. The interfacial affinity of CoAP accounts for the excellent catalytic activity observed by measuring the PCET current. Indeed, it is much more efficient than cobalt porphine and 2,3,7,8,12,13,17,18-octaethylporphyrin cobalt(II) (CoOEP) which are not amphiphilic (see Supporting Information, Figure S-3).33,35 The comparison with CoOEP highlights in particular the role of the amphiphlicity of the catalyst as both CoAP and CoOEP have the same redox potential for the first oxidation (0.69 V vs SHE).

Conclusions

In summary, the catalytic effect of (2,8,13,17-tetraethyl-3,7,12,18-tetramethyl-5-p-amino-phenylporphyrin) cobalt(II) (CoAP) on the proton-coupled oxygen reduction by Fc was studied at a water/DCE interface. CoAP serves as a redox catalyst like conventional monomeric cobalt porphyrins, activating O₂ for the reduction to H₂O₂. On the other hand, since CoAP has a strong interfacial affinity due to the amino group at a meso position, it presents an excellent electrocatalytic activity over other cobalt porphyrins that do not adsorb at the water/DCE interface such as the nonamphiphilic analogue CoOEP. This study demonstrates the concept of interfacial molecular electrocatalysis for the reduction of oxygen combining the use of molecular catalysts under electrochemical control.

In a general manner, the advantage of using amphiphilic molecular catalysts at polarized liquid—liquid interfaces stems from the fact that catalysts may be oxidized and destroyed during the course of the reaction by spurious radical side reactions. The degraded catalysts are likely to lose their amphiphility and to desorb from the interface where new catalysts will be able to adsorb from the bulk. This is similar to what happens in natural photosystems, in which the molecular catalysts have a finite lifetime and are replaced permanently by freshly synthesized ones.

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Supporting Information Available: Porphyrin synthesis procedures, measurements of redox potentials of CoAP, math derivation of surface excess concentration (eq 5), spectrophometric titration of CoAP with TFA, and the comparison of catalysis of CoAP with other cobalt porphyrins. This material is available free of charge via the Internet at http://pubs.acs.org.

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Figure 7. CVs in the presence of both 5 mM Fc and 25 μM CoAP (x = 25 μM, y = 5 mM) using Cell 1 at various pH, scan rate 0.02 V s⁻¹.

Figure 8. (a) CVs in the presence of 25 μM CoAP and 5 mM Fc (black), DFC (red), and FcCA (blue) using Cell 1 (x = 25 μM, y = 5 mM), scan rate 0.02 V s⁻¹; (b) Logarithmic plots of the catalytic current in the forward scan for Fc (black) and DFC (red).